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(54) Title: UNIVERSAL CORONAVIRUS VACCINE

### (57) Abstract

A universal vaccine is disclosed which elicits a protective immune response in different host species and against different coronaviruses. A polypeptide which elicits protective antibodies against a homologous sequence found in the C terminal portion of coronavirus S proteins is disclosed. Vaccines comprising either the polypeptide or nucleic acids which encode the polypeptide are also disclosed. Methods of protecting a host against coronavirus infection are disclosed.

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## Universal Coronavirus Vaccine

## Cross reference to related applications

This application is a continuation-in-part application of U.S. application serial number 07/882,171, filed May 8, 1992, pending, which is a continuation-in-part of U.S. application serial number 07/698,927, filed May 13, 1991, which is a continuation-in-part of U.S. application serial number 07/613,066, filed November 14, 1990, each of which is incorporated herein by reference.

#### 10 Field of the invention

The present invention relates to a universal vaccine useful to protect different species of animals against infection by different host-specific coronaviruses.

# Background of the invention

15 Coronaviruses are a family of host-specific enveloped RNA viruses with a single-stranded positive sense genome. Examples of coronaviruses include, but are not limited to: feline infectious peritonitis (FIPV) and feline enteric coronavirus (FECV) which are specific to felines; 20 canine coronavirus (CCV) which is specific to canines; transmissible gastroenteritis coronavirus (TGEV) which is specific to swine; bovine coronavirus (BCV) which is specific to bovine species; human coronavirus which is specific to humans; mouse hepatitis virus (MHV) which is specific to 25 murine species; and infectious bronchitis virus (IBV) which specific to avian species. These host-specific coronaviruses cannot cross infect different species of Viral infection of the host by a coronavirus can cause symptoms ranging from mild enteritis to severe 30 debilating disease to, in some cases, death.

Coronaviruses share common structural features including a spike or S protein (also referred to as a peplomer protein). The S protein is a glycoprotein which protrudes

from the surface of the virus particle. The S protein mediates the binding of virions to the host cell receptor and is involved in membrane fusion. In addition, it is the target of virus neutralizing antibodies.

5 S proteins contain an N-terminal signal sequence, a C-terminal transmembrane segment and potential N-linked glycosylation sites. Comparison of different coronavirus S proteins show little homology, i.e. similarity, at the N terminus and highly conserved amino acid sequences at the C 10 terminus. Because the tissue tropism and symptomatology is quite varied among this virus family, it is speculated that the pathogenesis of coronaviruses is determined by the sequences encoded at the N-terminus while the more conserved C-terminus encodes critical structural 15 features common to all coronaviruses. The carboxy terminus of the S protein is believed to be involved in fusion.

The structure of the S protein has been studied. Cavanagh (1983) J. Gen. Virol. 64:2577-2583, which is incorporated herein by reference, proposed a model for the coronavirus spike in which the C-terminal half of the protein forms its stalk and the N-terminal half, its bulbous protein. deGroot et al., (1987) J. Mol. Biol. 197:, which is incorporated herein by reference, have postulated a model in which a coiled-coil structure forms the connection between the globular part of the S protein and the viral membrane. This model is based on the occurrence of heptad repeats, i.e., a periodicity (a-b-c-d-e-f-g) in which the amino acids are hydrophobic. Britton (1991) Nature 353:394, which is incorporated herein by reference, reported the presence of a 30 leucine zipper motif at the carboxyl end of the S glycoprotein of coronaviruses for which the spike sequence is available: TGEV FS772/70 (amino acids 1342-1377), FIPV WSU 1146 (amino acids 1345-1380), MHV A59 (amino acids 1217-1252), human coronavirus 229E (amino acids 1067-1102), BCV Mebus (amino 35 acids 1266-1294), and infectious bronchitis virus Beaudette (amino acids 1059-1079). The leucine zipper motif terminates

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ten residues upstream of the conserved KWP motif preceding the transmembrane domain.

Efforts have been made to develop vaccines against various host-specific coronaviruses. Attempts have been made 5 with varying success to develop attenuated live virus vaccines, inactivated vaccines, subunit vaccines recombinant nucleic acid based vaccines. In each case, the vaccine developed did not cross-protect other host animals. currently available for protection 10 coronavirus are specific for protection against a given member of the coronavirus family. Such vaccines do not provide cross protection to protect a host against other members of the coronavirus family which are able to infect the species. Furthermore, such vaccines do not cross protect other animals 15 against coronaviruses for which they are susceptible to infection.

There is a need for a vaccine which can protect against coronavirus infection. In particular, there is a need for a vaccine which can be useful to protect a host species against different coronaviruses and there is a need for a vaccine which can be useful to protect different host species against different coronaviruses.

### Summary of the invention

The present invention relates to a polypeptide comprising an amino acid sequence from the C terminal portion of a coronavirus S protein which has been found to be highly conserved among coronaviruses and which is capable of eliciting a protective immune response. This sequence is referred to as a universal conserved domain. The polypeptides of the present invention have less than a complete amino acid sequence of an S protein.

The present invention relates to a vaccine comprising a polypeptide which includes an universal conserved domain and which has less than a complete amino acid sequence of an S protein.

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The present invention relates to an isolated nucleic acid molecule having a nucleic acid sequence which encodes a polypeptide that includes a universal conserved domain polypeptide and that has less than a complete amino acid 5 sequence of an S protein.

The present invention relates to a vaccine comprising a nucleic acid molecule that encodes a polypeptide which includes an universal conserved domain and which has less than a complete amino acid sequence of an S protein.

The present invention relates to a method of protecting an animal from infection by a coronavirus comprising administering an amount of a polypeptide effective to elicit a protective immune response. The polypeptide administered in the method comprises a universal conserved 15 domain and has less than a complete amino acid sequence of an S protein.

The present invention relates to a method of protecting an animal from infection by a coronavirus comprising administering an amount of a nucleic acid molecule 20 which encodes a polypeptide effective to elicit a protective immune response. The polypeptide encoded by the nucleic acid molecule administered in the method comprises a universal conserved domain and has less than a complete amino acid sequence of an S protein.

## 25 Detailed description of the invention

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According to the present invention, a highly conserved region of the spike protein has been identified which, when presented as a vaccine component or product, is useful as a universal immunogen to protect an animal against 30 coronavirus infection. The vaccine of the present invention may be used to vaccinate any animal susceptible to infection by virus that is a member of the coronavirus family. Accordingly, the present invention provides vaccines which can be produced in a single manufacturing process and administered 35 to different species of animals. The cross-protection afforded by vaccines of the present invention eliminates the

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need to produce different vaccines to protect animals against different members of the coronavirus family.

As used herein, the term "polypeptide" is meant to refer to a peptide, polypeptide or protein molecule; a molecule which includes a peptide, polypeptide or protein molecule; or a molecule that contains amino acid residues which are linked by non-peptide bonds.

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As used herein, the term "universal conserved domain" ("UCD") is meant to refer to the identical 124 amino 10 acid segment found in the C terminal portion of S proteins from TGEV, CCV and strains of feline coronaviruses. addition, the term "UCD" is meant to refer to the corresponding amino acid segments of other coronavirus which have different but homologous amino acid sequences. 15 corresponding sequences may be identified by their location in the S protein, i.e. downstream of the bulbous N-terminal region and upstream of the transmembrane region and the high level of amino acid sequence similarity to the 124 amino acid sequence described above. Furthermore, the term "UCD" is 20 additionally meant to refer to consensus sequences are generated by comparing corresponding sequences and determining the statistically average amino acid residue at a given position in the sequence. Thus, when several different sequences are compared, the most common residue at a given position is assigned to that position in a consensus sequence.

The conservation of UCD sequences suggests that they play a major role in virus structure and/or replication. The region of perfect homology decreases in size as other coronavirus S genes are included in the comparison. For example, bovine and human coronavirus are more closely aligned to the feline, canine and porcine coronavirus S genes in this conserved region than are sequences from the murine and avian coronaviruses.

Table 1 contains a comparison of corresponding amino 35 acid sequences from the C terminal portion of various coronaviruses. SEQ ID NO:1 is an amino acid sequence from FIPV strain Wsue2 (Virulent, Type II; Genbank accession number

X06170). SEQ ID NO:2 is an amino acid sequence from FIPV strain Df2e2 (Virulent, Type II). SEQ ID NO:3 is an amino acid sequence from FIPV strain Tse2 (Temperature sensitive mutant of Df2). SEQ ID NO:4 is an amino acid sequence from 5 FECV strain Fecve2 (Avirulent strain 1683). SEQ ID NO:5 is an amino acid sequence from TGEV strain Tgeve2 (Purdue strain; Genbank accession number D00118). SEQ ID NO:6 is an amino acid sequence from FIPV strain Tgeve2f2 (Miller strain; Genbank accession number M56002). SEQ ID NO:7 is an amino 10 acid sequence from BCV strain Bcve2 (Genbank accession number M30613). SEQ ID NO:8 is an amino acid sequence from HCV strain Hcve2 (Genbank accession number X16816). SEQ ID NO:9 is an amino acid sequence from IBV strain Ibbspi (Genbank accession number X16816). SEQ ID NO:10 is an amino acid 15 sequence from MHV strain Mhve2a59 (Genbank accession number X51939 SEQ ID NO:11 is an amino acid sequence from FIPV strain Mhvs (Genbank accession number X04797). SEQ ID NO:12 is a consensus sequence which has been designed to provide an optimum UCD amino acid sequence.

The 124 residue amino acid sequence which is completely conserved in TGEV, CCV and feline coronaviruses is shown in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4 and SEQ ID NO:5 from residue 37 to residue 160. The consensus sequence, SEQ ID NO:12, also contains this 124 amino acid 25 sequence in its entirety from residue 37 to residue 160. This 124 amino acid sequence is currently a preferred UCD sequence of the present invention. The entire 199 amino acid consensus sequence is a preferred UCD-containing peptide.

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Using amino acid sequence information from any 30 coronavirus, one having ordinary skill in the art can identify the conserved region corresponding to the 124 amino acid sequence found in TGEV, CCV and feline coronaviruses. exemplified in Table 1, the amino acid sequences from the C terminal portion of coronaviruses can be compared to identify 35 the sequence which corresponds to the UCD from TGEV, CCV and feline coronaviruses. The procedure is straightforward and

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can be performed to provide additional UCD sequences and flanking sequences.

corresponding conserved regions from coronaviruses other than CCV, TGEV and feline coronaviruses may be identified by their location on the S protein and the high level of sequence homology the possess when compared to the 124 amino acid sequence referred to above. An example of such comparison and identification is shown in Table 1 in which sequences from the C terminal regions of various S proteins upstream from the transmembrane region are compared and homologous sequences identified. Widely available computer programs such as PLOTSIMILARITY software (Genetics Computer Group, Madison WI) may be employed to locate a UCD in a coronavirus.

15 In addition, such software may be employed to expedite the generation of consensus sequences. This software relies on the principles originally set out by Wilbur and Lipman and later refined by Smith and Waterman and by Needleman and Wunsch. Using these well known guidelines, 20 having ordinary skill in the art may compare sequences and arrive at the statistically average or most common residue occupying a given position. The PLOTSIMILARITY software automates this function. Consensus sequences are thus generated. In addition to the consensus sequence provided as 25 SEQ ID NO:12, a different consensus sequence derived from a comparison of corresponding sequences is disclosed in the coowned, co-pending patent application: which is filed on the same day as the present application; which is entitled "Compositions and Methods for Vaccinating Coronaviruses"; 30 which names the same inventors as the present application (Miller, Timothy J.; Jones, Elaine V.; Reed, Albert P.; and Klepfer, Sharon R); which has been designated docket number H85009-1 by Applicants; and which is incorporated herein by reference.

Accordingly, the present invention relates to polypeptides which comprise a UCD or a fragment or a derivative thereof. That is, the present invention relates

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to polypeptides which comprise: the 124 amino acid sequence form TGEV, CCV and feline coronaviruses; or the different amino acid sequences from other coronaviruses which correspond to the 124 amino acid sequence; or a consensus sequence 5 generated from comparison of corresponding regions; or immunogenic fragments or immunogenic derivatives thereof.

Polypeptides according to the present may further comprise additional flanking sequences from coronavirus or flanking sequences designed as a consensus sequence of the flanking sequences of corresponding regions from different coronaviruses.

As used herein, the term "immunogenic fragment" is meant to refer to polypeptides which include an incomplete UCD which is capable of eliciting a protective immune response against coronavirus in an animal susceptible to coronavirus infection. Immunogenic fragments may comprise a sequence having nine or more amino acids from a UCD, and may include additional amino acid sequences.

is meant to refer to molecules which have a UCD or portions thereof with conservative amino acid substitutions and which are capable of eliciting a protective immune response against a coronavirus in an animal susceptible to coronavirus infection. Those having ordinary skill in the art can readily design derivatives having UCD sequences with conservative substitutions for amino acids. For example, following what are referred to as Dayhof's rules for amino acid substitution (Dayhof, M.D. (1978) Nat. Biomed. Res. Found., Washington, D.C. Vol. 5, supp. 3), amino acid residues in a peptide sequence may be substituted with comparable amino acid residues. Such substitutions are well known and are based the upon charge and structural characteristics of each amino acid.

Using standard procedures and readily available starting materials, one having ordinary skill in the art can determine whether a fragment and derivative is an immunogenic fragment or an immunogenic derivative, respectively. Briefly, polypeptides can be produced by standard methodologies and

tested to determine whether they are capable of eliciting a protective immune response. Sera from vaccinated animals can be analyzed to detect the presence of antibodies capable of inhibiting infection of cells in culture. Furthermore, 5 challenge studies can be performed to determine if animals vaccinated with a polypeptide are protected from subsequent infection by wild type virus. One having ordinary skill in the art can routinely produce and screen fragments and derivatives to determine the effectiveness of such vaccine 10 components to elicit protective immune responses. Similarly, larger molecules may also be screened by the same means to detect their ability to elicit a protective immune response.

The UCD lies near the transmembrane region of the S protein. Because this region of the S protein is purported 15 to be involved in the secondary structure of the glycoprotein, in receptor binding and in virus-induced cell fusion, the UCD plays an important role in the function of the S protein and in the formation of infectious virus. Inducing an immune response against this region will interfere with the folding 20 of the S glycoprotein into its proper conformation. presence of circulating antibodies to this region could bind to either virus or infected cells expressing the glycoprotein on the surface. Virus complexed with antibody may be unable to bind to receptors on susceptible cells and/or initiate the 25 pathway required to gain entry which involves a conformational change of the S protein. Recognition of this region on the surface of infected cells would target them for clearance. Antibody binding to the conserved region of the S protein surface expressed by infected cells would, most likely, prevent cell fusion and interfere with virus assembly. Regardless of mechanism, an immune response to the UCD of a coronavirus S protein will inhibit virus spread from cell to cell and limit virus infection.

Polypeptides according to the present invention 35 comprise less than a complete S protein sequence. particular, the polypeptides do not comprise a complete Nterminal portion of an S protein and preferably comprise few

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or no amino acid sequences from the N-terminal bulbous portion of the protein. Furthermore, the polypeptides preferably do not comprise a complete transmembrane domain of an S protein. In some preferred embodiments, polypeptides comprise no more than a 400 amino acid sequence upstream (from the C terminus to the N terminus) from about 2 amino acids upstream from the transmembrane domain. In some preferred embodiments, polypeptides comprise no more than a 300 amino acid sequence upstream (from the C terminus to the N terminus) from about 5 amino acids upstream from the transmembrane domain.

In some preferred embodiments, polypeptides which comprise a UCD, or derivatives and/or fragments thereof further comprise flanking sequences of the UCD found in coronavirus. For example, in some preferred embodiments, the polypeptide comprises portions of the S protein flanked by and optionally including the heptad repeats reported by deGroot et al., such as, for example, in FIPV strain WSU 1146 from residues 1067 to 1380. In some preferred embodiments, the polypeptide comprises portions of the S protein flanked on the carboxy side by and may also include a leucine zipper motif as reported by Britton. In some preferred embodiments, the polypeptide comprises portions of the S protein from about 300 residues upstream of the transmembrane region to about 5 amino acid residues upstream from the transmembrane domain.

In some preferred embodiments, the polypeptide comprises a UCD about 124 amino acids in length. In some preferred embodiments, the polypeptide comprises an immunogenic fragment of a UCD about 100 amino acids in length. In some preferred embodiments, the polypeptide comprises an immunogenic fragment of a UCD about 50 amino acids in length. In some preferred embodiments, the polypeptide comprises an immunogenic fragment of a UCD about 25 amino acids in length. In some preferred embodiments, the polypeptide comprises an immunogenic fragment of a UCD about 15 amino acids in length.

35 In some preferred embodiments, the polypeptide comprises an immunogenic fragment of a UCD about 10 amino acids in length.

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In some preferred embodiments, a UCD comprises amino acid residues 37-160 of SEQ ID NO:12. Additional preferred embodiments comprise SEQ ID NO:12. Other preferred embodiments of the invention comprise SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4 or SEQ ID NO:5. Other preferred embodiments comprise SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10 or SEQ ID NO:11.

In addition to a UCD and, optionally, additional flanking segments from an S protein, other peptide segments

10 may also be included in the polypeptide of the present invention. Such additional peptide segments may comprise other immunogenic targets from coronavirus and/or other pathogens, and/or they may be provided for improved stability, UCD epitope presentation or production/purification facilitation. The resulting polypeptide is considered a chimeric or fusion polypeptides.

Vaccines according to the present invention can be employed vaccinate animals against infection coronaviruses or at least to prevent the clinical symptoms 20 associated with such infections. Such vaccines will provide protection against multiple coronaviruses and cross species protection. Vaccines may be produced which are either protein-based or nucleic acid-based. In both cases, the vaccinated animal is exposed to an immunogenic polypeptide 25 which comprises a UCD. A protective immune response is elicited which is sufficient to protect the animal against coronavirus.

Vaccines according to the present invention can be either:

- a) compositions which comprise a polypeptide that includes a universal conserved domain; or
- b) compositions which comprise a nucleic acid molecule that includes a nucleotide sequence which encodes a polypeptide that includes a universal conserved domain. In
   35 both types of vaccines, the polypeptide is not a complete S protein and it elicits a protective immune response in animals.

In protein based, i.e. subunit vaccines, polypeptides having a UCD may by produced using standard techniques including recombinant DNA techniques for protein production or by peptide synthesis. In preferred embodiments, polypeptides used in subunit vaccines according to the present invention are produced by recombinant DNA methodology.

The nucleic acid sequences of coronavirus S genes are widely known. One having ordinary skill in the art may routinely obtain DNA that encodes a polypeptide including a 10 UCD using standard techniques and widely available starting materials. The nucleotide and amino acid sequences for S proteins from several types and strains of coronaviruses can found in the co-owned published PCT application PCT/US91/08525 which claims priority to U.S. Application Serial Numbers 613,066 and 698,927; each of these applications are incorporated herein by reference. Nucleotide and amino acid sequences of S proteins can also be found in published European Patent Applications publication numbers: 0,524,672 A1; 0,411,684 A2; 0,264,979 A1; 0,138,242 A1; and 20 application number EP 91 30 3737. Each of these European patent applications are incorporated herein by reference. In addition, nucleotide and amino acid sequences of S proteins from several coronaviruses as well as nucleotide and amino acid sequences of a consensus sequence is disclosed in the co-25 owned, co-pending patent application: which is filed on the same day as the present application; which is entitled "Compositions and Methods for Vaccinating Coronaviruses"; which names the same inventors as the present application (Miller, Timothy J.; Jones, Elaine V.; Reed, Albert P.; and 30 Klepfer, Sharon R); which has been designated docket number H85009-1 by Applicants; and which is incorporated herein by reference.

Nucleic acid molecules encoding some or all of an S protein from a coronavirus may be generated by a variety of techniques. For such molecules, a nucleotide sequence that encodes a UCD may be identified. Using, for example, Polymerase Chain Reaction (PCR) methodology, primers flanking

both sides the region of interest may be designed and used to produce multiple copies of the UCD routinely. Alternatively, using restriction enzymes, a UCD may be isolated from DNA encoding an S protein. Moreover, nucleic acid molecules that 5 encode a UCD may also be synthesized using techniques well known to those having ordinary skill in the art.

One having ordinary skill in the art can, using well techniques, insert such DNA molecules commercially available expression vector for use in well known 10 expression systems. For example, the commercially available plasmid pSE420 (Invitrogen, San Diego, CA) may be used for production of a DNA encoding a polypeptide including a UCD in coli. The commercially available plasmid pYES2 (Invitrogen, San Diego, CA) may, for example, be used for 15 production in S. cerevisiae strains of yeast. The commercially available MaxBac™ (Invitrogen, San Diego, CA) complete baculovirus expression system may, for example, be used for production in insect cells. The commercially available plasmid pcDNA I (Invitrogen, San Diego, CA) may, for 20 example, be used for production in mammalian cells such as Chinese Hamster Ovary cells. One having ordinary skill in the art can use these commercial expression vectors and systems or others to produce a polypeptide including a UCD using routine techniques and readily available starting materials. 25 (See e.g., Sambrook et al., Molecular Cloning a Laboratory Manual, Second Ed. Cold Spring Harbor Press (1989) which is incorporated herein by reference.) Thus, the desired proteins can be prepared in both prokaryotic and eukaryotic systems, resulting in a spectrum of processed forms of the protein.

The particulars for the construction of expression systems suitable for desired hosts are known to those in the art. Briefly, for recombinant production of the protein, the DNA encoding the polypeptide is suitably ligated into the expression vector of choice. The DNA is operably linked to 35 all regulatory elements which are necessary for expression of the DNA in the selected host. One having ordinary skill in

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the art can, using well known techniques, prepare expression vectors for recombinant production of the polypeptide.

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The expression vector including the DNA that encodes the polypeptide comprising a UCD is used to transform the 5 compatible host which is then cultured and maintained under conditions wherein expression of the foreign DNA takes place. The protein of the present invention thus produced is recovered from the culture, either by lysing the cells or from the culture medium as appropriate and known to those in the 10 art. One having ordinary skill in the art can, using well known techniques, isolate the polypeptide that includes a UCD produced using such expression systems.

In addition to producing these proteins recombinant techniques, automated peptide synthesizers may 15 also be employed to produce polypeptides that include a UCD. Such techniques are well known to those having ordinary skill in the art and are useful if derivatives which have substitutions not provided for in DNA-encoded protein production.

Subunit vaccines according to the invention comprise a polypeptide the includes a UCD but which is not a complete S protein and a pharmaceutically acceptable carrier or diluent. Optionally, the vaccine may comprise additional immunogenic proteins, additional vaccine components such as 25 non-subunit vaccines, and/or an adjuvant.

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In nucleic acid molecule based, i.e. recombinant vaccines, a nucleotide sequences which encode polypeptides that include a UCD is inserted into a vector and administered to the animal. The vector delivers genetic material to the 30 animal where it is transcribed and translated to produce the immunogenic polypeptide. Vectors for use as vaccines are well known and include non-pathogenic viruses and prokaryotic organisms. Suitable vectors for delivering genetic material are readily available or may be produced from readily 35 available starting materials using standard techniques. Two examples of vectors useful for delivering genetic material as a vaccine are the recombinant pox vectors or non-pathogenic

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Salmonella strains. The nucleotide sequence that encodes the immunogenic polypeptide is operably linked to regulatory elements required for expression and inserted within the vector. Alternatively, it is incorporated into the vector at 5 a site where it is placed under the control of the necessary regulatory elements already present in the vector. Naked DNA may also be used as a vaccine delivery system.

Recombinant vaccines may be used in combination with other vaccines. Further, the genetic material which encodes 10 the polypeptide that comprises the UCD may further comprise additional coding sequences which encode other peptide sequences capable of eliciting an immunogenic response against coronavirus or another pathogen.

Both subunit and recombinant vaccines may 15 formulated following accepted convention using buffers, stabilizers, preservative, solubilizers and compositions used to facilitate sustained release. Generally, additives for isotonicity can include sodium chloride, dextrose, mannitol, sorbitol and lactose. Stabilizers include gelatin and 20 albumin. Adjuvants such as aluminum or magnesium hydroxide may be employed. Vaccines may be maintained in solution or, in some cases, particularly recombinant vaccines, lyophilized. Lyophilized vaccine may be stored conveniently and combined with sterile solution before administration.

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The amount of polypeptide administered depends upon such factors as the size of the polypeptide, the species, age, weight, and general physical characteristics of the animal, and by the composition of the vaccine. Determination of optimum dosage for each parameter may be made by routine 30 methods. Generally, subunit vaccines according to the present invention contain between 0.05-5000 micrograms of polypeptide per milliliter of sterile solution, preferably 10-1000 micrograms. Generally, recombinant vaccines according to the present invention contain between 105-108 infectious units per 35 milliliter of sterile solution. About .5-2 milliliter of polypeptide-containing solution is administered.

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Subunit vaccines and genetic material based vaccines may be administered by an appropriate route such as, for example, by oral, intranasal, intramuscular, intraperitoneal or subcutaneous administration. In some embodiments, 5 intranasal or subcutaneous administration is preferred. Subsequent to initial vaccination, animals may be boosted by revaccination.

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### Examples

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Example 1 Cloning of Coronavirus Conserved Region in pMG1

The bacterial expression vector, pMG-1, allows a gene expressing a foreign protein to be fused to a partial sequence of the NS1 gene from influenza virus, the first 81 encoding amino acids thereof. This vector is described in European Patent Application No. 366,238, published May 2, 15 1990, which is incorporated herein by reference.

Primers were designed to amplify a S gene region encoding amino acids 1115-1238 of the DF2 FIPV strain for expression in this vector as follows. The upstream primer contains NcoI and NdeI restriction sites and initiates 20 amplification at base pair 3406 (amino acid 1115), and is SEQ ID NO:13:

5'-GTTGTCAACACACCATGGATCATATGCAAGGGCAAGCTTTAAGTCACCTTACA. <u>Nco</u>I NdeI

25 The downstream primer contains a StuI site and terminates amplification at base pair 3777 (amino acid 1238), and is SEQ ID NO: 14:

5'-AAATACCTGAGGCCTCCAAGCTGTTACAGTTTCATAAGCTGT. StuI

30 The amplified fragment (412 bp) was cloned into the pT, Blue vector according to the manufacturer's instructions. plasmid containing amino acids 1115-1238 in pT7 Blue was digested with NcoI/StuI, the 412 base pair insert isolated, and ligated overnight at 15°C to plasmid vector pMG1 digested 35 with NcoI/StuI and dephosphorylated. Host cells AR120 and AR58 were transformed with the ligation mix and the presence

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of insert bearing clones was confirmed by diagnostic restriction enzyme digestions.

Vaccinia recombinants were engineered to contain the 1115-1238 amino acid conserved region of WT DF2 FIPV. The conserved region was cloned into the vaccinia expression vector pSC11 by blunt-ending the 412 base pairs NcoI/StuI fragment isolated from the pT7 Blue clone described in Example 12, end-filling by incubation with Klenow polymerase, and inserting it into the SmaI site downstream of the 7.5K vaccinia promoter. The ligation mix was transformed into HB101 host cells. Full-length clones were identified and oriented with respect to vector by BamHI and ScaI digests of mini-prep DNAs, respectively.

#### Table 1

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	Wsue2	NITQAFGKVN	DAIHQTSQGL	ATVAKALAKV	<b>QDVVNTQGQA</b>	LSHLTVQLQN
	Df2e2	NITQAFGKVN	DAIHQTSQGL	ATVAKALAKV	<b>QDVVNTQGQA</b>	LSHLTVQLQN
5	Tse2	NITQAFGKVN	DAIHQTSQGL	ATVAKALAKV	<b>QDVVNTQQQA</b>	LSHLTVQLQN
	Fecve2			ATVAKALAKV		
	Tgeve2	NITOAFGKVN	DAIHOTSOGL	ATVAKALAKV	ODVVNTOGOA	LSHLTVOLON
	Tgeve2f2	NITOAFGKVN	DAIHOTSOGL	ATVAKALAKV	ODVVNTOGOA	LSHLTVOLON
	Bcve2	AIOEGFDATN	S	ALVKI	CAVVNANARA	LNNTLOOLSN
10	Hcve2	NIVDAFTGVN	DATTOTSOAT.	QTVATALNKI	ODVVNOOGNS	LNHTTSOLRO
	Ibbspi	HMOR.	GP	RSTSLALQQI	ODVVSKOSAT	T.TRTWAST.NK
	Mhve2a59			ALGKI		
	Mhys	ATORGRDATN	9	ALGKI	VOLLIMIY MY BY	TNNTTNOTCH
		NITQAFGKVN	DATHOTS GL	ATVAVALAKV	ODVINITAGO	T CHT TUOT CM
		WII OWA	mitudio: gr	UT AUTOMUTA	δη α αυτ δαδυ	TOUT LAST ON
15		51			•	100
	Wsue2		DTYNRIDELS	ADAQVDRLIT	CDT.TAT.NARU	
	Df2e2	NEOATSSSIS	DIVNRIDELS	ADAQVDRLIT	CDITALNARY	COLLINGUEA
	Tse2			ADAQVDRLIT		
	Fecve2			ADAOVDRLIT		
20	Tgeve2			ADAQVDRLIT		
20	Tgeve2f2			ADAQVDRLIT		
	Bcve2			AQAQIDRLIN		
	Hcve2			ADQQVDRLIT		
	Ibbspi			ANAQVDRLIT		
25	Mhve2a59	DECUTERATO	PIOGGEDATA	AKAQIDRLIN	CULMALMANA	PAVÕAPOTKA
	Mhvs	PREATERSTO	ETT TOT DAVID	AKAQIDRLIN	CULTATIONALI	SVATSDSITT
		NFQAISSSIS	PILITADAYE	SUNCTION THE	CULTATIVALIT	SVOTSDSIFT
		WIANIDOOLD	DITHAMDEDS	MDVÖADVETT	GREINERAFV	POITTY NAME A
		101		•		
		TOT				150
	Wsue2		VNECVRSQSQ	RFGFCGNGTH	LFSLANAAPN	
30	Wsue2 Df2e2	RASRQLAKDK	VNECVRSQSQ VNECVRSQSQ	RFGFCGNGTH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN	GMIFFHTVLL
30		RASRQLAKDK RASRQLAKDK	VNECVRSQSQ	RFGFCGNGTH	LFSLANAAPN	GMIFFHTVLL GMIFFHTVLL
30	Df2e2	RASRQLAKDK RASRQLAKDK RASRQLAKDK	VNECVRSQSQ VNECVRSQSQ	RFGFCGNGTH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL
30	Df2e2 Tse2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ	RFGFCGNGTH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL
30	Df2e2 Tse2 Fecve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL
30 35	Df2e2 Tse2 Fecve2 Tgeve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL
	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV
	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSS VNECVKSQSK	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL
	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVKSQSS VNECVKSQSK INECVKSQSI	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPY IFSIVNAAPE VLTIPQNAPN	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYPIHFSYV GLVFLHTVLL GIVFIHFSYT
	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK KVSAAQAIEK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSK VNECVKSQSK INECVKSQSI VNECVKSQSI	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGHH RYGFCGNGTH RYSFCGNGRH RYSFCGNGRH RINFCGNGNH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GIVFIHFSYT GLYFIHFSYV
35	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KVSAAQAIEK KFSAAQAIEK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSS VNECVKSQSS VNECVKSQSI VNECVKSQTT VNECVKSQTT	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGTH RINFCGNGNH RINFCGNGNH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVONAPY	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYPIHFSYV GLVFLHTVLL GLYPIHFSYV GLYPIHFSYV GLYPIHFSYV
	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK KVSAAQAIEK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSS VNECVKSQSS VNECVKSQSI VNECVKSQTT VNECVKSQTT	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGTH RINFCGNGNH RINFCGNGNH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVONAPY	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYPIHFSYV GLVFLHTVLL GLYPIHFSYV GLYPIHFSYV GLYPIHFSYV
35	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KVSAAQAIEK KFSAAQAIEK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSS VNECVKSQSS VNECVKSQSI VNECVKSQTT VNECVKSQTT	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGTH RINFCGNGNH RINFCGNGNH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVONAPY	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYPIHFSYV GLVFLHTVLL GLYPIHFSYV GLYPIHFSYV GLYPIHFSYV
35	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK RASRQLAKDK	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSK INECVKSQST VNECVKSQTT VNECVRSQST VNECVRSQSQ	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGHH RINFCGNGNH RINFCGNGNH RINFCGNGNH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVQNAPY LFSLANAAPN	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLYFIHFSYT GLYFIHFSYV GLYFIHFSYV GLYFIHFSYV GMIFFHTVLL
35	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK 151 PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSS VNECVKSQST VNECVKSQTT VNECVKSQTT VNECVRSQSQ SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH RINFCGNGNH RINFCGNGNH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVQNAPY LFSLANAAPN LTSLANAAPN LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYT GLYFIHFSYV GLCFIHFSYV GMIFFHTVLL
35	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK 151 PTAYETVTAW PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVRSQSK INECVRSQST VNECVRSQTT VNECVRSQTT VNECVRSQSQ SGICASDGDR SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH RINFCGNGNH RINFCGNGNH RINFCGNGNH RFGFCGNGTH	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYT GLYFIHFSYV GLCFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ FYLTPRTMYQ
35	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK  151 PTAYETVTAW PTAYETVTAW PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVRSQSK INECVRSQST VNECVRSQTT VNECVRSQTT VNECVRSQTT VNECVRSQSQ SGICASDGDR SGICASDGDR SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH RINFCGNGNH RINFCGNGNH RINFCGNGNH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK LTLFRNLDDK LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYT GLYFIHFSYV GLCFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ
<b>35</b>	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS Wsue2 Df2e2 Tse2 Fecve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQQRELATQK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK  151 PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVRSQSK INECVRSQST VNECVRSQTT VNECVRSQTT VNECVRSQTT VNECVRSQSQ SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH RINFCGNGNH RINFCGNGNH RINFCGNGNH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK LTLFRNLDDK LTLFRNLDDK LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYV GLYFIHFSYV GLYFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ
<b>35</b>	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS Wsue2 Df2e2 Tse2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQQK SQORELATQK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK  151 PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVKSQSI VNECVKSQSI VNECVKSQTT VNECVKSQTT VNECVRSQSQ SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RYSFCGNGNH RYSFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPY IISLVQNAPY ILSLVQNAPY ILSLVQNAPY LFSLANAAPN LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYV GLYFIHFSYV GLYFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ
<b>35</b>	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS Wsue2 Df2e2 Tse2 Fecve2 Tgeve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATQK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK  151 PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVKSQSS VNECVKSQSI VNECVKSQSI VNECVKSQTT VNECVKSQTT VNECVRSQSQ SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH RFGFCGNGTH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY LFSLANAAPN LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYV GLYFIHFSYV GLYFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ FYLTPRTMYQ
<b>35</b>	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS Wsue2 Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2 Tgeve2f2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KVSAAQAIEK KFSAAQAIEK RASRQLAKDK  151 PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVKSQSS VNECVKSQSI VNECVKSQSI VNECVKSQTT VNECVKSQTT VNECVRSQSQ SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK SGYFVNVNNT	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYFIHFSYV GLVFLHTVLL GLYFIHFSYV GLYFIHFSYV GLYFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ WMFTGSGYYY
35 40 45	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2 Bcve2 Hcve2 Hbspi Mhve2a59 Mhv8 CONSENSUS Wsue2 Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2 Tgeve2 Ecve2 Hcve2 Hcve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KFSAAQAIEK KFSAAQAIEK RASRQLAKDK  151 PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSSQ VNECVRSQSS VNECVKSQSI VNECVKSQSI VNECVKSQTT VNECVKSQTT VNECVRSQTO SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH RINFCGNGNH RINFCGNGNH RINFCGNGNH RFGFCGNGTH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYPIHFSYV GLVPIHFSYV GLVPIHFSYV GLYPIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ
<b>35</b>	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS Wsue2 Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2 Tgeve2f2 Bcve2	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAYDK SQQRELATOK KVSAAQAIEK RASRQLAKDK 151 PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVKSQSS VNECVKSQSS VNECVKSQST VNECVKSQTT VNECVKSQTT VNECVRSQSQ SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RYSFCGNGRH RINFCGNGNH RINFCGNGNH RINFCGNGNH RFGFCGNGTH TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK LTLFRNLDCK	GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GMIFFHTVLL GLYPIHFSYV GLVPIHFSYV GLVPIHFSYV GLCFIHFSYV GMIFFHTVLL  200 FYLTPRTMYQ WMFTGSGYYY YRITSRIMFE YYITARDMYM
35 40 45	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Hbspi Mhve2a59 Mhvs CONSENSUS Wsue2 Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Lbspi	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KVSAAQAIEK RASRQLAKDK 151 PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSSQ VNECVKSQSS VNECVKSQSS VNECVKSQST VNECVKSQTT VNECVKSQTT VNECVRSQTT VNECVRSQSQ SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYGFCGNGTH RINFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH RFGFCGNGTH TFGLVVKDVQ	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPE VLTIPQNAPN ILSLVQNAPY ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK LTLFRNLDK LT	GMIFFHTVLL  200 FYLTPRTMYQ WMFTGSGYYY YRITSRIMFE YYITARDMYM WKFTGSSYYY
35 40 45	Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs CONSENSUS  Wsue2 Df2e2 Tse2 Fecve2 Tgeve2 Tgeve2f2 Bcve2 Hcve2 Ibbspi Mhve2a59 Mhvs	RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK RASRQLAKDK KFSAAQAMEK RASRQLAQOK SQQRELATOK KVSAAQAIEK RASRQLAKDK 151 PTAYETVTAW	VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSQ VNECVRSQSS VNECVRSQSS VNECVRSQST VNECVRSQST VNECVRSQST VNECVRSQST VNECVRSQTT VNECVRSQTT VNECVRSQTT VNECVRSQSQ SGICASDGDR	RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RFGFCGNGTH RINFCGNGNH RYSFCGNGTH RINFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH RINFCGNGNH TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ TFGLVVKDVQ GIAPK TNGYVLRQPN SQUAIVPANG GLAPK GLAPK	LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN LFSLANAAPN IISLVQNAPY IFSIVNAAPS VLTIPQNAPY ILSLVQNAPY LFSLANAAPN  LTLFRNLDDK LTLFRNLDK LTLFRNLDDK LTLFRNLDDK LTLFRNLDK LTLFRNL	GMIFFHTVLL  200 FYLTPRTMYQ WMFTGSGYYY YRITSRIMFE YYITARDMYM WKFTGSSYYY WKFTGSNYYY

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#### SEQUENCE LISTING

(1) GENERAL INFORMATION: (i) APPLICANT: Miller, Timothy J. Jones, Blaine V. 5 Reed, Albert P. Klepfer, Sharon R. (ii) TITLE OF INVENTION: Universal Coronavirus Vaccine (iii) NUMBER OF SEQUENCES: 14 10 (iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: SmithKline Beecham Corporation (B) STREET: 709 Swedeland Road (C) CITY: King of Prussia (D) STATE: PA 15 (E) COUNTRY: USA (F) ZIP: 19406-2799 (V) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible 20 (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: 25 (C) CLASSIFICATION: (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 07/882,171 (B) FILING DATE: 08-MAY-1992 (vii) PRIOR APPLICATION DATA: 30 (A) APPLICATION NUMBER: US 07/698,927 (B) FILING DATE: 13-MAY-1991 (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 07/613,066 (B) FILING DATE: 14-NOV-1990 35 (viii) ATTORNEY/AGENT INFORMATION: (A) NAME: Schreck, Patricia A. (B) REGISTRATION NUMBER: 33,777 (C) REFERENCE/DOCKET NUMBER: SBC/PAS/WW001 (2) INFORMATION FOR SEQ ID NO:1: 40 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 200 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein 45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1: Asn Ile Thr Gln Ala Phe Gly Lys Val Asn Asp Ala Ile His Gln Thr

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•	Ser	Gln	Gly	Leu 20	Ala	Thr	Val	Ala	Lys 25	Ala	Leu	Ala	Lys	Val 30	Gln	Asp
	Val	Val	Asn 35	Thr	Gln	Gly	Gln	Ala 40	Leu	Ser	His	Leu	Thr 45	Val	Gln	Leu
5	Gln	Asn 50	Asn	Phe	Gln	Ala	Ile 55	Ser	Ser	Ser	Ile	Ser 60	Asp	Ile	Tyr	Asn
	Arg 65	Leu	Asp	Glu	Leu	Ser 70	Ala	Asp	Ala	Gln	Val 75	Asp	Arg	Leu	Ile	Thr 80
10	Gly	Arg	Leu	Thr	Ala 85	Leu	Asn	Ala	Phe	Val 90	Ser	Gln	Thr	Leu	Thr 95	Arg
	Gln	Ala	Glu	Val 100	Arg	Ala	Ser	Arg	Gln 105	Leu	Ala	ГÀв	Asp	Lys 110	Val	Asn
	Glu	Сув	Val 115	Arg	Ser	Gln	Ser	Gln 120	Arg	Phe	Gly	Phe	Cys 125	Gly	Asn	Gly
15	Thr	His 130	Leu	Phe	Ser	Leu	Ala 135	Asn	Ala	Ala	Pro	Asn 140	Gly	Met	Ile	Phe
	Phe 145	His	Thr	Val	Leu	Leu 150	Pro	The	Ala	Tyr	Glu 155	Thr	Val	Thr	Ala	Trp 160
20	Ser	Gly	Ile	Сув	Ala 165	Ser	Asp	Gly	Asp	Arg 170	Thr	Phe	Gly	Leu	Val 175	Val
	Lys	Asp	Val	Gln 180	Leu	Thr	Leu	Phe	Arg 185	Asn	Leu	Asp	Asp	Lys 190	Phe	Tyr
	Leu	Thr	Pro 195	Arg	Thr	Met	Tyr	Gln 200			-					
25	(2) INFO	RMAT	ON 1	OR S	SEQ :	ED NO	):2:									
	. (i)	(B)	JENCI LEI TYI	NGTH:	200 amin	am:	ino a id		3							
30	(ii)	MOL	3CUL	3 TY	PE: ]	prote	≘in				-					
	(xi)	SEQ	DENC	S DES	SCRI	PTIO	N: S	BQ. II	ONO	:2:						
	Asn 1	Ile	Thr	Gln	Ala 5	Phe	Gly	Lys	Val	Asn 10	Asp	Ala	Ile	His	Gln 15	Thr
35	Ser	Gln	Gly	Leu 20	Ala	Thr	Val	Ala	Lys 25	Ala	Leu	Ala	Lys	Val 30	Gln	Asp
	Val	Val	Asn 35	Thr	Gln	Gly	Gln	Ala 40	Leu	Ser	His	Leu	Thr 45	Val	Gln	Leu
	Gln	Asn 50	Asn	Phe	Gln	Ala	Ile 55	Ser	Ser	Ser	Ile	Ser 60	Asp	Ile	Tyr	Asn
40	Arg 65	Leu	yab	Glu	Leu	Ser 70	Ala	Asp	Ala	Gln	Val 75	Asp	Arg	Ĺeu	Ile	Thr 80
	Gly	Arg	Leu	Thr	Ala 85	Leu	Asn	Ala	Phe	Val 90	Ser	Gln	Thr	Leu	Thr 95	Arg

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Gln Ala Glu Val Arg Ala Ser Arg Gln Leu Ala Lys Asp Lys Val Asn. 105 Glu Cys Val Arg Ser Gln Ser Gln Arg Phe Gly Phe Cys Gly Asn Gly 115 5 Thr His Leu Phe Ser Leu Ala Asn Ala Ala Pro Asn Gly Met Ile Phe Phe His Thr Val Leu Leu Pro Thr Ala Tyr Glu Thr Val Thr Ala Trp Ser Gly Ile Cys Ala Ser Asp Gly Asp Arg Thr Phe Gly Leu Val Val 10 Lys Asp Val Gln Leu Thr Leu Phe Arg Asn Leu Asp Asp Lys Phe Tyr 185 Leu Thr Pro Arg Thr Met Tyr Gln 195 15 (2) INFORMATION FOR SEQ ID NO: 3: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 200 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear 20 (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3: Asn Ile Thr Gln Ala Phe Gly Lys Val Asn Asp Ala Ile His Gln Thr Ser Gln Gly Leu Ala Thr Val Ala Lys Ala Leu Ala Lys Val Gln Asp 25 Val Val Asn Thr Gln Gly Gln Ala Leu Ser His Leu Thr Val Gln Leu Gln Asn Asn Phe Gln Ala Ile Ser Ser Ser Ile Ser Asp Ile Tyr Asn 30 Arg Leu Asp Glu Leu Ser Ala Asp Ala Gln Val Asp Arg Leu Ile Thr Gly Arg Leu Thr Ala Leu Asn Ala Phe Val Ser Gln Thr Leu Thr Arg Gln Ala Glu Val Arg Ala Ser Arg Gln Leu Ala Lys Asp Lys Val Asn 35 105 Glu Cys Val Arg Ser Gln Ser Gln Arg Phe Gly Phe Cys Gly Asn Gly Thr His Leu Phe Ser Leu Ala Asn Ala Ala Pro Asn Gly Met Ile Phe 40 Phe His Thr Val Leu Leu Pro Thr Ala Tyr Glu Thr Val Thr Ala Trp 150 Ser Gly Ile Cys Ala Ser Asp Gly Asp Arg Thr Phe Gly Leu Val Val

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Lys Asp Val Gln Leu Thr Leu Phe Arg Asn Leu Asp Asp Lys Phe Tyr 180 185 190

Leu Thr Pro Arg Thr Met Tyr Gln 195 200

- 5 (2) INFORMATION FOR SEQ ID NO:4:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 200 amino acids
    - (B) TYPE: amino acid(D) TOPOLOGY: linear
- 10 (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Asn Ile Thr Gln Ala Phe Gly Lys Val Asn Asp Ala Ile His Gln Thr 1 5 10 15

Ser Gln Gly Leu Ala Thr Val Ala Lys Ala Leu Ala Lys Val Gln Asp 20 25 30

Val Val Asn Thr Gln Gly Gln Ala Leu Ser His Leu Thr Val Gln Leu 35 40 45

Gln Asn Asn Phe Gln Ala Ile Ser Ser Ser Ile Ser Asp Ile Tyr Asn 50 55 60

20 Arg Leu Asp Glu Leu Ser Ala Asp Ala Gln Val Asp Arg Leu Ile Thr 65 70 75 80

Gly Arg Leu Thr Ala Leu Asn Ala Phe Val Ser Gln Thr Leu Thr Arg 85 90 95

Gln Ala Glu Val Arg Ala Ser Arg Gln Leu Ala Lys Asp Lys Val Asn 100 105 110

Glu Cys Val Arg Ser Gln Ser Gln Arg Phe Gly Phe Cys Gly Asn Gly 115 120 125

Thr His Leu Phe Ser Leu Ala Asn Ala Ala Pro Asn Gly Met Ile Phe 130 140

Phe His Thr Val Leu Leu Pro Thr Ala Tyr Glu Thr Val Thr Ala Trp
145 150 155 160

Ser Gly Ile Cys Ala Ser Asp Gly Asp Arg Thr Phe Gly Leu Val Val 165 170 175

Lys Asp Val Gln Leu Thr Leu Phe Arg Asn Leu Asp Asp Lys Phe Tyr
180 185 190

Leu Thr Pro Arg Thr Met Tyr Gln 195 200

(2) INFORMATION FOR SEQ ID NO:5:

40

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 200 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

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		(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	ои о	:5:						
		Asn 1	Ile	Thr	Gln	Ala 5	Phe	Gly	Lys	Val	Asn 10	Asp	Àla	Ile	His	Gln 15	Thr
5		Ser	Gln	Gly	Leu 20	Ala	Thr	Val	Ala	Lys 25	Ala	Leu	Ala	Lys	Val 30	Gln	Asp
		Val	Val	Asn 35	Thr	Gln	Gly	Gln	Ala 40	Leu	Ser	His	Leu	Thr 45	Val	Gln	Leu
		Gln	Asn 50	Asn	Phe	Gln	Ala	Ile 55	Ser	Ser	Ser	Ile	Ser 60	yab	Ile	Tyr	Asn
10		Arg 65	Leu	Asp	Glu	Leu	Ser 70	Ala	Asp	Ala	Gln	Val 75	Asp	Arg	Leu	Ile	Thr 80
		Gly	Arg	Leu	Thr	Ala 85	Leu	Asn	Ala	Phe	Val 90	Ser	Gln	Thr	Leu	Thr 95	Arg
15		Gln	Ala	Glu	Val 100	Arg	Ala	Ser	Arg	Gln 105	Leu	Ala	Lys	Asp	Lys 110	Val	Asn
		Glu	Cys	Val 115	Arg	Ser	Gln	Ser	Gln 120	Arg	Phe	Gly	Phe	Cys 125	Gly	Asn	Gly
		Thr	His 130	Leu	Phe	Ser	Leu	Ala 135	Asn	Ala	Ala	Pro	Asn 140	Gly	Met	Ile	Phe
20		Phe 145	His	Thr	Val	Leu	Leu 150	Pro	Thr	Ala	Tyr	Glu 155	Thr	Val	Thr	Ala	Trp 160
		Ser	Gly	Ile	Сув	Ala 165	Ser	Asp	Gly	Asp	Arg 170	Thr	Phe	Gly	Leu	Val 175	Val
25		ГÀв	yab	Val	Gln 180	Leu	Thr	Leu	Phe	<b>Arg</b> 185	Asn	Leu	Asp	Asp	Lys 190	Phe	Tyr <sub>.</sub>
		Leu	Thr	Pro 195	Arg	Thr	Met	Tyr	Gln 200								
	(2)	INFOI	[TAM	ON F	OR S	EQ I	D NO	:6:									
30		(i)	(A) (B)	LEN	GTH: E: a	RACT 200 mino Y: 1	ami aci	TICS no a d	:: icids	3							
	(	(ii)	MOLE	CULE	TYP	E: p	rote	in									
	(	(xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	6:						
35		Asn 1	lle	Thr	Gln	Ala 5	Phe	Gly	Lys	Val	Asn 10	Asp	Ala	Ile	His	Gln 15	Thr
		Ser	Gln	Gly	Leu 20	Ala	Thr	Val	Ala	Lys 25	Ala	Leu	Ala	Lys	Val 30	Gln	Asp
40		Val	Val	Asn 35	Thr	Gln	Gly	Gln	Ala 40	Leu	Ser	His	Leu	Thr 45	Val	Gln	Leu
		Gln	Asn 50	Asn	Phe	Gln	Ala	Ile 55	Ser	Ser	Ser	Ile	Ser 60	Asp	Ile	Tyr	Asn

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							•	- 24	<b>!</b> -		-				•	
	Arg 65	Leu	Asp	Glu	Leu	Ser 70	Ala	Asp	Ala	Gln	Val 75	Asp	Arg	Leu	Ile	Thr 80
	Gly	Arg	Leu	Thr	Ala 85	Leu	Asn	Ala	Phe	Val 90	Ser	Gln	Thr	Leu	Thr 95	Arg
5	Gln	Ala	Glu	<b>Val</b> 100	Arg	Ala	Ser	Arg	Gln 105	Leu	Ala	ГÄв	Авр	Lув 110	Val	Asn
	Glu	Cys	Val 115	Arg	Ser	Gln	Ser	Gln 120	Arg	Phe	Gly	Phe	Cys 125	Gly	Asn	Gly
10	Thr	His 130	Leu	Phe	Ser	Leu	Ala 135	Asn	Ala	Ala	Pro	Asn 140	Gly	Met	Ile	Phe
	Phe 145	His	Thr	Val	Leu	Leu 150	Pro	Thr	Ala	Tyr	Glu 155	Thr	Val	Thr	Ala	Trp 160
	Ser	Gly	Ile	Cya	Ala 165		Asp	Gly	Asp	Arg 170	Thr	Phe	Gly	Leu	Val 175	Val
15	ГÀв	Asp	Val	Gln 180	Leu	Thr	Leu	Phe	Arg 185	Asn	Leu	Asp	Asp	Lys 190	Phe	Tyr
	Leu	Thr	Pro 195	Arg	Thr	Met	Tyr	Gln 200				•	•			
	(2) INFO	RMAT:	ION 1	FOR S	SEQ 1	D NO	0:7:									
20	<b>(i)</b>	(B	JENCI LEI TYI	NGTH:	179 mino	ami	ino a id		3							
	(ii)	MOLI														
25	(xi)	SEQ	JENCI	, DES	CRIE	TIO	N: SI	g II	) NO:	:7:						
	Ala 1	Ile	Gln	Glu	Gly 5	Phe	Авр	Ala	Thr	Asn 10	Ser	Ala	Leu	Val	Lys 15	Ile
	Gln	Ala	Val	Val 20	Asn	Ala	Asn	Ala	Glu 25	Ala	Leu	Asn	Asn	Leu 30	Leu	Gln
30	Gln	Leu	Ser 35	Asn	Arg	Phe	Gly	Ala 40	Ile	Ser	Ser	Ser	Leu 45	Gln	Glu	Ile .
	Leu	Ser 50	Arg	Leu	yab	Ala	Leu 55	Glu	Ala	Gln	Ala	Gln 60	Ile	Asp	Arg	Leu
35	Ile 65	Asn	Gly	Arg	Leu	Thr 70	Ala	Leu	Asn	Val	Tyr 75	Val	Ser	Gln	Gln	Leu 80
	Ser	yab	Ser	Thr	Leu 85	Val	Lys	Phe	Ser	Ala 90	Ala	Gln	Ala	Met	Glu 95	Lys
	Val	Asn	Glu	Cys 100	Val	Lys	Ser	Gln	Ser 105	Ser	Arg	Ile	Asn	Phe 110	Gly	Asn
40	Gly	Asn	His 115	Ile	Ile	Ser	Leu	Val 120	Gln	Asn	Ala	Pro ·	Tyr 125	Gly	Leu	Tyr
	Phe	Ile 130	His	Phe	Ser	Tyr	Val 135	Pro	Thr	Lys	Tyr	Val 140	Thr	Ala	Lys	Tyr

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Ser Pro Gly Leu Cys Ile Ala Gly Asp Arg Gly Ile Ala Pro Lys Ser-

Gly Tyr Phe Val Asn Val Asn Asn Thr Trp Met Phe Thr Gly Ser Gly

5 Tyr Tyr Tyr

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- (2) INFORMATION FOR SEQ ID NO:8:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 196 amino acids(B) TYPE: amino acid
- 10
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
- Asn Ile Val Asp Ala Phe Thr Gly Val Asn Asp Ala Ile Thr Gln Thr 15

Ser Gln Ala Leu Gln Thr Val Ala Thr Ala Leu Asn Lys Ile Gln Asp

Val Val Asn Gln Gln Gly Asn Ser Leu Asn His Leu Thr Ser Gln Leu

20 Arg Gln Asn Phe Gln Ala Ile Ser Ser Ser Ile Gln Ala Ile Tyr Asp

Arg Leu Asp Thr Ile Gln Ala Asp Gln Gln Val Asp Arg Leu Ile Thr

Gly Arg Leu Ala Ala Leu Asn Val Phe Val Ser His Thr Leu Thr Lys 25

Tyr Thr Glu Val Arg Ala Ser Arg Gln Leu Ala Gln Gln Lys Val Asn

Glu Cys Val Lys Ser Gln Ser Lys Arg Tyr Gly Phe Cys Gly Asn Gly

30 Thr His Ile Phe Ser Ile Val Asn Ala Ala Pro Glu Gly Leu Val Phe 135

> Leu His Thr Val Leu Leu Pro Thr Gln Tyr Lys Asp Val Glu Ala Trp 150

Ser Gly Leu Cys Val Asp Gly Thr Asn Gly Tyr Val Leu Arg Gln Pro 35

> Asn Leu Ala Leu Tyr Lys Glu Gly Asn Tyr Tyr Arg Ile Thr Ser Arg 180

Ile Met Phe Glu 195

- 40 (2) INFORMATION FOR SEQ ID NO:9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 183 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear

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(ii)	MOLECULE	TYPE:	protein
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	(xi)	SEQ	JENCI	B DES	SCRI	PTIO	N: SI	EQ II	NO:	:9:						
	His 1	Met	Gln	Glu	Gly 5	Phe	Arg	Ser	Thr	Ser 10	Leu	Ala	Leu	Gln	Gln 15	Ile
5	Gln	Asp	Val	Val. 20	Ser	Гла	Gln	Ser	Ala 25	Ile	Leu	Thr	Glu	Thr 30	Met	Ala
	Ser	Leu	Asn 35	Lys	Asn	Phe	Gly	Ala 40	Ile	Ser	Ser	Val	Ile <b>4</b> 5	Gln	Glu	Ile
10	Gln	Gln 50	Phe	yab	Ala	Ile	G1n 55	Ala	Asn	Ala	Gln	Val 60	yeb	Arg	Leu	Ile
	Thr 65	Gly	Arg	Leu	Ser	Ser 70	Leu	Ser	Val	Leu	Ala 75	Ser	Ala	Lys	Gln	Ala 80
	Glu	Ile	Arg	Val	Ser 85	Gln	Gln	Arg	Glu	Leu 90	Ala	Thr	Gln	Lys	Ile 95	Ası
15	Glu	Сув	<b>Val</b>	Lys 100	Ser	Gln	Ser	Ile	Arg 105	Tyr	Ser	Phe	Cys	Gly 110	Asn	Gl
	Arg	His	Val 115	Leu	Thr	Ile	Pro	Gln 120	Asn	Ala	Pro	Asn	Gly 125	Ile	Val	Phe
20	Ile	His 130	Phe	Ser	Tyr	Thr	Pro 135	Asp	Ser	Phe	Val	Asn 140	Val	Thr	Ala	Ile
	Val 145	Gly	Phe	Cys	Val	Lys 150	Pro	Ala	Asn	Ala	Ser 155	Gln	Ala	Ile	Val	Pro 160
	Ala	Asn	Gly	Arg	Gly 165	Ile	Phe	Ile	Gln	Val 170	Asn	Gly	Ser	Tyr	Tyr 175	Ile
25	Thr	Ala	Arq	Asp	Met	Tvr	Met									

- 5 Thr Ala Arg Asp Met Tyr Met 180
  - (2) INFORMATION FOR SEQ ID NO:10:
    - (i) SEQUENCE CHARACTERISTICS:
      (A) LENGTH: 180 amino acids
  - (B) TYPE: amino acid

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- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Ala Ile Glm Asp Gly Phe Asp Ala Thr Asn Ser Ala Leu Gly Lys Ile
  1 5 10 15
  - Gln Ser Val Val Asn Ala Asn Ala Glu Ala Leu Asn Asn Leu Leu Asn 20 25 30
  - Gln Leu Ser Asn Arg Phe Gly Ala Ile Ser Ala Ser Leu Gln Glu Ile 35 40 45
- 40 Leu Thr Arg Leu Glu Ala Val Glu Ala Lys Ala Gln Ile Asp Arg Leu
  50 55

- 27 -

Ile Asn Gly Arg Leu Thr Ala Leu Asn Ala Tyr Ile Ser Lys Gln Leu . 65 70 75 80 Ser Asp Ser Thr Leu Ile Lys Val Ser Ala Ala Gln Ala Ile Glu Lys 5 Val Asn Glu Cys Val Lys Ser Gln Thr Thr Arg Ile Asn Phe Cys Gly Asn Gly Asn His Ile Leu Ser Leu Val Gln Asn Ala Pro Tyr Gly Leu Tyr Phe Ile His Phe Ser Tyr Val Pro Ile Ser Phe Thr Thr Ala Asn 10 Val Ser Pro Gly Leu Cys Ile Ser Gly Asp Arg Gly Leu Ala Pro Lys Ala Gly Tyr Phe Val Gln Asp Asp Gly Glu Trp Lys Phe Thr Gly Ser 15 Ser Tyr Tyr Tyr (2) INFORMATION FOR SEQ ID NO:11: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 180 amino acids 20 (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11: Ala Ile Gln Glu Gly Phe Asp Ala Thr Asn Ser Ala Leu Gly Lys Ile 25 Gln Ser Val Val Asn Ala Asn Ala Glu Ala Leu Asn Asn Leu Leu Asn Gln Leu Ser Asn Arg Phe Gly Ala Ile Ser Ala Ser Leu Gln Glu Ile 30 Leu Thr Arg Leu Asp Ala Val Glu Ala Lys Ala Gln Ile Asp Arg Leu Ile Asn Gly Arg Leu Thr Ala Leu Asn Ala Tyr Ile Ser Lys Gln Leu Ser Asp Ser Thr Leu Ile Lys Phe Ser Ala Ala Gln Ala Ile Glu Lys 35 Val Asn Glu Cys Val Lys Ser Gln Thr Thr Arg Ile Asn Phe Cys Gly Asn Gly Asn His Ile Leu Ser Leu Val Gln Asn Ala Pro Tyr Gly Leu 40 Cys Phe Ile His Phe Ser Tyr Val Pro Thr Ser Phe Lys Thr Ala Asn Val Ser Pro Gly Leu Cys Ile Ser Gly Asp Arg Gly Leu Ala Pro Lys 145

Ala Gly Tyr Phe Val Gln Asp Asn Gly Glu Trp Lys Phe Thr Gly Ser 165 170 175

Asn Tyr Tyr Tyr 180

- 5 (2) INFORMATION FOR SEQ ID NO:12:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 199 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
- 10 (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Asn Ile Thr Gln Ala Phe Gly Lys Val Asn Asp Ala Ile His Gln Thr 1 5 10 15

Ser Gly Leu Ala Thr Val Ala Lys Ala Leu Ala Lys Val Gln Asp Val
15 25 30

Val Asn Thr Gln Gly Gln Ala Leu Ser His Leu Thr Val Gln Leu Gly 35 40 45

Asn Asn Phe Gln Ala Ile Ser Ser Ser Ile Ser Asp Ile Tyr Asn Arg 50 55 60

20 · Leu Asp Glu Leu Ser Ala Asp Ala Gln Val Asp Arg Leu Île Thr Gly 65 70 75 80

Arg Leu Thr Ala Leu Asn Ala Phe Val Ser Gln Thr Leu Thr Arg Gln 85 90 95

Ala Glu Val Arg Ala Ser Arg Gln Leu Ala Lys Asp Lys Val Asn Glu

25 100 105 110

Cys Val Arg Ser Gln Ser Gln Arg Phe Gly Phe Cys Gly Asn Gly Thr 115 120 125

His Leu Phe Ser Leu Ala Asn Ala Pro Asn Gly Met Ile Phe Phe 130 140

30 His Thr Val Leu Leu Pro Thr Ala Tyr Glu Thr Val Thr Ala Trp Pro 145 150 155 160

Gly Ile Cys Ala Ser Asp Gly Asp Arg Thr Phe Gly Leu Val Val Lys 165 170 175

Asp Val Gln Leu Thr Leu Phe Arg Asn Leu Asp Asp Lys Phe Tyr Leu 35 180 185 190

Thr Pro Arg Thr Met Tyr Gln 195

(2) INFORMATION FOR SEQ ID NO:13:

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- (i) SEQUENCE CHARACTERISTICS:
  (A) LENGTH: 53 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA

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		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
		GTTGTCAACA CACCATGGAT CATATGCAAG GGCAAGCTTT AAGTCACCTT ACA	53
•		(2) INFORMATION FOR SEQ ID NO:14:	
	5	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 42 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii) MOLECULE TYPE: cDNA	
	10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
		AAATACCTGA GGCCTCCAAG CTGTTACAGT TTCATAAGCT GT	42

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#### Claims

1. A polypeptide comprising a universal conserved domain of a coronavirus or an immunogenic fragment or derivative thereof; said polypeptide having less than a 5 complete amino acid sequence of said S protein.

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- A vaccine comprising a pharmaceutically acceptable carrier or diluent and a polypeptide comprising a universal conserved domain of a coronavirus or an immunogenic fragment or derivative thereof; said polypeptide having less than a
   complete amino acid sequence of said S protein.
- 3. A nucleic acid molecule comprising a nucleotide sequence that encodes a polypeptide comprising a universal conserved domain of a coronavirus or an immunogenic fragment or derivative thereof; said polypeptide having less than a complete amino acid sequence of said S protein.
- 4. A recombinant vaccine comprising a nucleic acid molecule, said nucleic acid molecule comprising a nucleotide sequence that encodes a polypeptide comprising a universal conserved domain of a coronavirus or an immunogenic fragment or derivative thereof; said polypeptide having less than a complete amino acid sequence of said S protein.
- 5. A method of protecting an animal against coronavirus comprising administering a polypeptide comprising a universal conserved domain of a coronavirus or an immunogenic fragment or derivative thereof; said polypeptide having less than a complete amino acid sequence of said S protein.
- 6. A method of protecting an animal against coronavirus comprising administering a nucleic acid molecule comprising a nucleotide sequence that encodes a polypeptide comprising 30 a universal conserved domain of a coronavirus or an immunogenic fragment or derivative thereof; said polypeptide

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having less than a complete amino acid sequence of said S protein.

## INTERNATIONAL SEARCH REPORT

Inumational application No.

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IPC(5) US CL	ASSIFICATION OF SUBJECT MATTER :C07K 3/00; C07H 15/12; C12N 15/00; A61K 39/1 :530/350; 536/27; 435/320.1; 424/89 to International Patent Classification (IPC) or to both							
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Minimum d	ocumentation searched (classification system followe	d by classification symbols)						
U.S. :	530/350; 536/27; 435/320.1; 424/89							
Documentat	tion searched other than minimum documentation to th	e extent that such documents are included	in the fields searched					
Electronic d	lata base consulted during the international search (na	ame of data base and, where practicable	, search terms used)					
	e Extra Sheet.	·	•					
C. DOC	UMENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.					
Y	EP, A, 0,264,979 (deGroot et al) document.	27 April 1988, see entire	1-6					
Y .	·							
X Furth	er documents are listed in the continuation of Box C	. See patent family annex.						
	cial categories of cited documents:	"I" inter document published after the inter date and not in conflict with the applica	tion but cited to understand the					
to b	ument defining the general state of the art which is not considered e part of particular relevance	principle or theory underlying the inve	ention					
	ier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be considered.	red to involve an inventive step					
cite	ument which may throw doubts on priority claim(a) or which is d to establish the publication date of another citation or other	when the document is taken alone  "Y"  document of particular columns the						
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'P" doc	ument published prior to the international filing date but later than priority date claimed	being obvious to a person skilled in th  *&" document member of the same patent						
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## INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/04365

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Y	The Journal of General Virology, Volume 71, No. 5, issued May 1990, T. Raabe et al., "Nucleotide Sequence of the Gene Encoding the Spike Glycoprotein of Human Coronavirus HCV	1-6
<b>Y</b>	229E", pp. 1065-1073, see entire document.  Archives of Virology, Volume 117, issued 1991, T. Hohdatsu et al., "Characterization of Monoclonal Antibodies Against Feline Infectious Peritonitis Virus Type II and Antigenic Relationship Between Feline, Porcine, and Canine Coronaviruses", pp. 85-95, see entire document.	1-6
Y	Virology, Volume 174, No. 2, issued February 1990, C. Sanchez et al., "Antigenic Homology Among Coronaviruses Related to Transmissable Gastroenteritis Virus", pp. 410-417, see entire document.	1-6

## INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/04365

B. FIELDS SEARCHED  Electronic data bases consulted (Name of data base and where practicable terms used):	
EMBL, GenBank, GeneSeq, PIR, Swiss-Prot, CA, Biosis, Medline, Embase, WPI, APS search terms: coronavirus, conserv?, spike, peplomer, C-term?, vaccine	
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